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Research Article

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Efficient Method of Lignin Isolation Using Microwave-Assisted Acidolysis and Characterization of the Residual Lignin

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- 7 Supporting Information

ABSTRACT: Microwave heating is characterized by high efficiency and selectivity in biomass treatment. Due to the high thermal stability and low polarity of lignin, isolation of lignin by high-temperature microwave treatment is a promising subject for investigation. In this paper, microwave treatment is applied to polysaccharide liquefaction and lignin isolation from softwood at 160–210 °C for 10 min with dilute sulfuric acid. Mass balance/element analysis/FTIR/TG/solid-state ¹³C NMR/Py-GC/MS are applied to investigate the processed residues (residual lignin). At 190 °C processing temperature, the residual lignin is a material rich in



aromatics. High lignin purity (93 wt %) and yield (82 wt %) could be achieved by a simple protocol, which usually takes days or even weeks using conventional milled wood lignin protocols. The Py-GC/MS is applied to check the structure of lignin by a newly developed approach. The liquid phase after isolation is analyzed by GC-MS and liquid carbon NMR. Most chemicals in processed liquid are from cellulose and hemicellulose, suggesting that lignin is preserved well in the residue. By comparison, we found that microwave isolation causes less lignin degradation than conventional acidolysis under equivalent conditions. It is concluded that microwave treatment is potentially a promising tool for isolation of polysaccharide-free lignin with high efficiency.

KEYWORDS: Lignin, Microwave, Acidolysis, Lignocellulosic biomass

↓ INTRODUCTION

25 Since 1838 when Anselme Payen first found "encrusting 26 material" that was later named "lignin" embedded between 27 cellulose and hemicellulose, numerous studies have been 28 carried out to investigate the structure and characteristics of 29 lignin. Lignin ranks second in quantity in the terrestrial regions 30 of Earth's surface, playing an important role in plants allowing 31 water conduction and protecting them against pathogen 32 attacks. From the viewpoint of chemical structure, lignin can 33 be a potential source of valuable phenolic compounds by 34 degradation. Compared with other sustainable carbon-based 35 resources, these vast resources constitute a potential advantage 36 for lignin utilization.

However, the extraction of polysaccharide-free lignin with high efficiency using conventional methods is still a challenge because in biomass lignin acts as "glue" adhering the plant polysaccharides layers together with strong covalent bonding to cellulose and hemicellulose. Extraction of lignin is accompanied by structural damage and polysaccharide contamination. The lack of high-quality lignin on the market coupled with difficulties in degrading it selectively and efficiently into useful low molecular weight products make it undervalued and underdeveloped compared with cellulose and hemicellulose. Therefore, lignin is still widely used as an energy source in chemical pulp and paper mills and in some industrial biorefinery processes. One advancement in pure lignin so isolation was proposed by Klason. The two-step Klason

acidolysis protocol and its modified versions have been mostly 51 used as standards of lignin content and purity determination, as 52 in the TAPPI T222 method.9 The drawback with the Klason 53 protocol is that as concentrated sulfuric acid is applied the 54 structure of Klason lignin (KL) is modified. In a KL procedure, 55 lignin condenses to become water-insoluble. As a result, the 56 repolymerization is serious. Another commonly used lignin in 57 laboratory studies is milled wood lignin (MWL). This milder 58 protocol uses neutral solvents for isolation affords and a 59 product that is widely regarded to offer the best material for the 60 structural analysis of the "native lignin" originally present in the 61 plant tissue. 5,10 The linkages in lignin—carbohydrate complexes 62 (LCC) are broken by milling, and then lignin is extracted by 63 dioxane-water solvent. The disadvantage is that the intensive 64 and lengthy milling (taking between 1 h to 3 weeks depending 65 on milling machine) is energy-consuming, which in turn 66 increases the cost of the isolation. Low lignin yield, 5 67 polysaccharide contamination, 11 and the tendency for diox- 68 ane-water to dissolve only the lower molecular weight 69 fractions of the lignin are also drawbacks of MWL protocol. 70 MWL is generally representative of total lignin in wood except 71 that phenolic content is higher than that in native lignin 72 because MWL is extracted by dioxane-water solvent. On the 73

Received: October 22, 2016 Revised: March 13, 2017 Published: March 30, 2017 74 basis of the MWL protocol, cellulolytic enzyme lignin (CEL) 75 protocol was proposed to increase lignin yield, but it was still 76 low at 27-29 wt %. 11 Furthermore, CEL protocol requires a 77 high dosage of enzymes, and the process is tedious. Therefore, 78 both MWL and CEL methods are used mainly by lab-scale 79 research but are not suitable for industrial production. 5,1 80 Although there are many improvements based on these 81 methods, efficient lignin isolation with high yield and low 82 contamination is always a difficult task and calls for new

With efficient and selective heating, microwave heating 85 provides a promising approach in thermal treatment of 86 biowaste, especially lignocellulose. 12-14 Until now, there have 87 been only a few studies focusing on microwave-assisted lignin 88 isolation. Zhou et al. 15 investigated microwave-assisted lignin 89 extraction from birch in formic acid and compared it with 90 conventional isolation methods. A higher delignification was 91 achieved by microwave heating than oil bath heating. Li et al. 16 92 also performed microwave lignin extraction from bamboo at 90 93 and 109 °C separately. It was found that increasing temperature 94 would benefit lignin extraction. Zoia et al. 17 performed 95 microwave-assisted lignin isolation in inorganic acid solution, 96 and a high yield of 55 wt % (total amount of acid soluble and 97 insoluble lignin) was achieved. All these studies prove the 98 advantages of microwave-assisted lignin isolation, especially 99 lignin purity and processing time. However, these studies only 100 focus on low-temperature isolation. High-temperature isolation 101 still needs investigation. With elevated temperature, better 102 performance is expected to be achieved because acidolysis 103 lignin is more stable to thermal degradation than cellulose and 104 hemicellulose. This thermal stability can be expected to be 105 further enhanced during microwave treatment because of the 106 selectivity of microwave treatment. Microwave heating is based on the high-frequency rotation of polar molecules. Therefore, 108 compounds with high polarity are more rapidly heated during 109 microwave irradiation. Lignin, having higher aromaticity and 110 lower polarity than polysaccharide, 18,19 is likely to degrade less 111 severely in a microwave isolation than conventional acidolysis 112 under equivalent conditions of total energy input.

Based on the discussion above, in this paper a new method 114 for fast microwave-assisted lignin isolation is proposed. Dilute 115 sulfuric acid is used for acidolysis, as previous studies have 116 shown that lignin-carbohydrate complexes (LCC) are reduced 117 to negligible levels when acidolysis is conducted in this 118 medium. Tigh-temperature isolation (160–210 °C) is carried 119 out to ensure LCC can be cleaved in a short time. Systematic 120 analysis is performed to investigate lignin quality. A new 121 analysis approach based on Py-GC/MS is applied to check the 122 structure of lignin after isolation.

MATERIALS AND METHODS 123

Materials. Mixed softwood pellets (MSP, UK Biochar Research 124 125 Centre, School of Geosciences, University of Edinburgh) were used as 126 feedstock for lignin isolation. The elemental and ICP analyses are 127 shown in Tables S1 and S2. Compared with hardwood and herbaceous 128 biomass, softwood has the least acid-soluble lignin, only about 0.2-0.5 129 wt %, and thus is the most suitable for acidolysis lignin isolation. 130 Sulfuric acid was purchased from Fischer Chemicals (>95 wt %). 131 Creosol (99 wt %), vanillin (99 wt %), and phenol, 2-methoxy- (98 wt 132 %) were purchased from Sigma-Aldrich. trans-Isoeugenol (99 wt %) 133 was purchased from Acros Organic.

Experimental Methods. All biomass was milled to 60 mesh 135 powders using a cutting mill (Retsch SM300, Germany) in 136 Biorenewables Development Center (BDC), University of York. The

microwave treatment was performed in a Discovery SP microwave 137 reactor (CEM Corporation, USA) in capped vessels. Maximum power 138 (300 W) of the microwave reactor was applied in all the experiments 139 to make sure that the holding temperature could be achieved as 140 quickly as possible. Diluted sulfuric acid (0.2 mol/L) was applied for 141 isolation. The processing temperature of 160-210 °C at intervals of 10 142 °C was used for isolation. The holding time was 5/10/20 min (in this 143 paper, the abbreviation microwave residual lignin (MRL) only refers to 144 the 10 min sample). During microwave treatment, 0.2 g of MSP and 145 15 mL of acid solvent were heated in a capped vessel with stirring. 146 After microwave treatment, the residue was recovered by filtration. 147 Then, the residue was washed several times with deionized water until 148 the rinsed water was neutral. In order to prepare the microwave 149 residual lignin obtained in this way for further analysis, the residue was 150 dried (105 °C, 24 h) and then weighed. All the experiments were 151 repeated 3 times.

Lignin isolation by conventional heating (acidolysis lignin, AL) was 153 performed using a benchtop autoclave (Anton Paar Monowave 50). 154 MSP (0.08 g) and aqueous sulfuric acid (0.2 mol/L, 6 mL) were 155 heated with stirring in a sealed vessel. The temperature was ramped up 156 to 190 °C (within 5 min, similar to microwave experiments) and was 157 held for 10 min. The residue (190 °C AL) after isolation was washed 158 and dried as in the microwave residual lignin preparation. Most 159 conditions of AL protocol are the same as those in the microwave 160 experiment. By comparing AL and MRL, the characteristics and 161 advantages of microwave treatment can be investigated.

The purity and yield was calculated by TAPPI T222 method. The 163 method is shown schematically in Figure S1. About 0.1 g of dewaxed 164 sample was treated with 10 g of sulfuric acid (72 wt %) at 20 °C for 2 165 h. The solution was then diluted with deionized water to 3 wt % 166 sulfuric acid and refluxed for 4h. The insoluble residue (lignin) was 167 isolated by filtration. After washed with hot water, the residue was 168 dried at 105 °C for 24 h. This dried residue is Klason lignin (KL). The 169 purity and yield were calculated according to the equation in Table 1. 170 tl The purity result was adjusted by subtracting the ash content 171 measured by TG analysis.

Table 1. Purity and Yield of MSP and 190 °C MRL/AL^a

	p		
	dry basis	extractive-free basis	yield (wt %)
MSP	30.37 ^b	39.08 ^c	
190 °C MRL	80.64 ^d	92.85 ^e	82.31 ^f
190 °C AL	75.91 ^d	87.51 ^e	65.60 ^f

^aFor definitions of M_0 , M_d , M_{al} , M_{i} , M_{di} , and M_{a2} , see Figure S1. $^bM_{a1}$ / M_0 . ${}^cM_{a1}/M_d$. ${}^dM_{a2}/M_i$. ${}^eM_{a2}/M_{di}$. ${}^fM_{a2}/M_{a1}$.

Elemental analysis and ICP analysis data were obtained from the 173 analytical service offered by Department of Chemistry, University of 174

Thermogravimetric (TG) analysis was performed using a Netzsch 176 STA 409 analyzer (Germany). The following parameters were applied: 177 temperature ramp rate 20 K/min, final temperature 600 °C, and carrier 178 gas 50 mL/min pure nitrogen gas. To measure ash content, the 179 following parameters were applied: temperature ramp rate 20 K/min, 180 final temperature 625 °C holding for 1 h, and carrier gas 50 mL/min 181 N₂ and 100 mL/min O₂. The final mass % was used as the ash content. 182

FTIR data was obtained using a PerkinElmer FTIR/FTNIR 183 Spectrum 400 analyzer (USA). The spectra were acquired between 184 700 and 4000 cm⁻¹ with resolution of 2 cm⁻¹ and scan time of 64 s. 185

Solid-state ¹³C NMR spectroscopy (SSNMR) results were obtained 186 at the EPSRC UK National Solid-State NMR Service at University of 187 Durham. The spectra were obtained at 100.562 MHz. The chemical 188 shift range from 0 to 240 ppm was recorded.

Py-GC/MS results were obtained from BDC, University of York. 190 The units used were CDS Analytical 5250-T Trapping Pyrolysis 191 Autosampler (UK) as the pyrolysis unit, Agilent Technologies 7890B 192 GC System (USA) as gas chromatography unit, and Agilent 193

194 Technologies 5977A MSD (USA) as mass spectrum unit. The sample 195 was loaded into the pyrolysis unit and pyrolyzed at 600 °C for 10 s. 196 The volatile materials released were carried into the GC/MS unit by 197 nitrogen for analysis. The following GC/MS parameters were applied: 198 GC inlet temperature at 350 °C, initial temperature at 40 °C for 2 min, 199 ramp rate at 10 K/min until 300 °C, holding at 300 °C for 30 min, and 200 split ratio with 50:1. Volatile compounds were identified by comparing 201 the mass spectra with NIST Lab database. A standard sample mixture 202 of four compounds, creosol/vanillin/2-methoxyphenol (guaiacol)/E-203 isoeugenol, was also subjected to pyrolysis and GC/MS in order to 204 verify the mass spectral identities.

After microwave isolation at 190 °C, the aqueous phase was neutralized and dried using freeze-dryer for 24 h, preparing for liquid-207 state 13 C NMR and GC/MS analysis. Liquid-state 13 C NMR 208 spectroscopy results were obtained by JEOL ECS 400 NMR 209 Spectrometer (Japan). D_2 O was used as the solvent for analysis. 210 The number of scans was 8192.

GC/MS results were obtained using a PerkinElmer Clarus 500 GC/212 MS (USA). Ethanol was chosen as solvent for analysis. The GC 213 program used was as follows: initial temperature at 50 °C holding for 4 214 min, ramp rate with 10 K/min until 290 °C and holding for 10 min, 215 split ratio with 5:1, and injector temperature at 290 °C. The identities 216 of the compounds were determined by comparing the mass spectra 217 with NIST lab database.

18 RESULTS AND DISCUSSION

Mass Balance and C/H Contents. Figure 1 shows the influence of temperature and holding time on the yield of

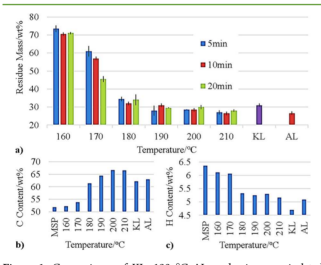


Figure 1. Comparisons of KL, 190 $^{\circ}$ C AL, and microwave-isolated lignin under different conditions: (a) mass balance; (b and c) C/H content.

221 residue and C/H content. It was found that major changes of 222 residue mass and C/H content took place from 160 to 190 °C. 223 At 170 °C, the residue mass could be still affected by holding 224 time; above 190 °C, the masses of the residues obtained did not 225 vary significantly with holding time, showing the high efficiency 226 of microwave heating. The residue mass of KL was 31 wt %, 227 which was close to that of 190/200 °C residues. Adler 228 reported that the equivalent formula of the purest softwood 229 lignin he could produce from spruce was 230 $C_9H_{7,92}O_{2.40}(OCH_3)_{0.92}$, which suggested the C/H content of 231 pure softwood lignin should be close to 65.12/5.47 wt %. Lin et 232 al. 22 also measured the C/H content of an industrial lignin 233 which was 65.00/6.43 wt %. Compared with these values, the 234 C/H content of 190 °C MRL was similar to those isolated 235 lignins. The H content of KL was low because concentrated

sulfuric acid was likely to dehydrate the lignin. The C/H of 190 236 °C AL was similar to that of 190 °C MRL; however, the residue 237 mass was lower, showing conventional acidolysis at high 238 temperature caused more mass loss than microwave treatment 239 and indicating that lignin is in fact more thermally stable under 240 microwave heating than when subjected to conventional 241 heating as expected.

Purities (Acid-Insoluble Lignin Content) and Yields. 243 The yields and purities of 190 °C MRL and 190 °C AL (10 min 244 sample) shown in Table 1 were calculated according to TAPPI 245 method T222 shown in Figure S1. After 10 min of microwave 246 treatment at 190 °C, the isolation produced lignin with high 247 purity (93 wt %) and yield (82 wt %), both of which were 248 higher than those of lignins obtained using MWL methods. Wu 249 and Argyropoulos¹¹ produced MWL with a 14 day milling 250 process on the softwood (black spruce (Picea mariana)). The 251 extractive-free basis purity and yields were 88.3 wt %/28.5 wt % 252 respectively. Compared to MWL methods, the much shorter 253 duration required by microwave heating is probably the main 254 reason for the higher lignin yields obtained during this study. 255 Within such a short processing time, lignin loss is reduced to a 256 great extent. Sulfuric acid offers an environment where 257 carbohydrate can be hydrolyzed and solubilized, while most 258 lignin is insoluble.

Another reason for high purity and yield is possibly the 260 selectivity of microwave treatment. 12–14 Different from conventional heating, microwave heating is achieved by the high-262 frequency rotation of polar molecules. Compared to nonpolar 263 compounds, polar molecules and functional groups are treated 264 more intensely and faster in microwave radiation. Compared 265 with carbohydrate, lignin is generally regarded as having higher 266 aromaticity and lower polarity. 18,19 Therefore, carbohydrate and 267 lignin can be expected to behave in significantly different ways 268 under microwave radiation, particularly in the presence of dilute 269 aqueous sulfuric acid. Such a hypothesis explains why in Table 270 1 the yield of 190 °C AL was much lower than that of 190 °C 271 MRL. These data provide further strong support for the 272 mechanism by which microwave heating exerts its selectivity in 273 mixtures containing materials of differing polarities.

Liquid-Phase Analysis. After isolation at 190 °C (10 min), 275 the solution after microwave treatment was analyzed by GC/ 276 MS and liquid ¹³C NMR. The GC/MS list of aqueous phase ²⁷⁷ compounds are showed in Table S3. The GC/MS results 278 showed that the majority of compounds in solution were 279 chemicals derived from sugars characterized by the presence of 280 ketone, aldehyde, and furan groups, while aromatic compounds 281 occurred in much lower proportions. This result was consistent 282 with liquid ¹³C NMR results (Figure S2). The peaks in 20-40 283 ppm were ascribed as saturated carbon which were mainly from 284 polysaccharide. The peaks of ketone were located in 205-220 285 ppm, suggesting that dehydrated sugars were probably the main 286 products in liquid phase. Of greatest significance is the absence 287 of intense peaks between 100 and 150 ppm, where carbons in 288 benzenoid rings typically resonate, indicating that aromatic 289 compounds remained predominantly in the insoluble solid 290 residue. The absence of these peaks provides further evidence 291 for lack of thermal depolymerization when lignin is heated by 292 microwaves for 10 min at 190 °C.

Thermogravimetric Analysis. Figure 2 shows the TG 294 f2 curves of MSP, KL, 170/190 °C MRL, and 190 °C AL. For 295 MSP, the DTG curve had a very strong peak at around 350 °C 296 that corresponds to the decomposition of cellulose. ^{23,24} This 297 peak was accompanied by a well-pronounced shoulder at 298

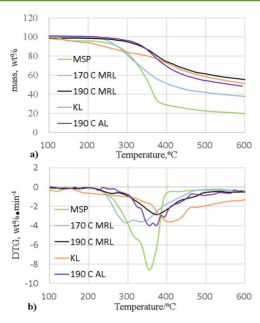


Figure 2. Pyrolysis curves of samples (MSP, KL 190 $^{\circ}$ C AL, and 170/190 $^{\circ}$ C MRL) at heating rate of 20 K/min. (a) TG curves; (b) DTG curves

299 around 300 °C, attributable to hemicellulose decomposi-300 tion. ^{23,24} For 170 °C MRL, the final mass loss was lower 301 than that of MSP. These results illustrated that microwave 302 isolation at 170 °C was already able to remove the carbohydrate 303 to some extent. However, there were two strong DTG peaks 304 (294 and 330 °C) in the range of 290–350 °C, showing that 305 170 °C MRL was still severely contaminated by cellulose and 306 hemicellulose.

The mass and DTG curves of KL and 190 °C MRL had 307 308 similar trends in general. Compared with linear structure of 309 cellulose and hemicellulose (with some branches), the complex 310 3D structure of lignin and predominance of aryl-alkyl ether 311 linkages make it recalcitrant to thermal degradation. These 312 factors resulted in the 190 °C MRL and KL samples having 313 high residual mass at 600 °C, a higher peak zone for 314 degradation. The final residual masses were high at 52 and 315 55 wt % respectively, showing that fewer degradable 316 compounds existed in these two samples than those in 170 °C MRL. Their DTG peaks were located between 370 and 410 318 °C, where pure lignin displays its DTG peak according to 319 previous studies. ^{23,24} Unlike the DTG curves of MSP and 170 °C MRL, the DTG curves showed no peaks between 290 to 321 350 °C, confirming that polysaccharides were mostly removed 322 in the 190 °C MRL and KL samples. A subtle difference between KL and 190 °C MRL was that the degrading peak of 324 190 °C MRL was slightly lower, which was either caused by structural changes brought about by the 190 °C treatment, or by dehydrations promoted by the 72 wt % sulfuric acid used in 327 the Klason protocol.

Comparing 190 °C MRL and 190 °C AL, it was found there was more polysaccharide in 190 °AL sample. The two DTG curves both had peaks at around 375 °C, showing lignin was a main component in both isolated residues. However, for the DTG curve of 190 °C AL, there was also a well-pronounced peak at 354 °C that was attributable to the degradation of polysaccharide. 23,24 Furthermore, the DTG peaks of 190 °C AL was less thermally stable. The data indicate that 190 °C AL was less thermally stable. The data indicate that 190 °C

MRL is less contaminated by polysaccharides and more 337 thermally stable than lignin produced by conventional acid- 338 olysis at 190 °C.

FTIR. Figure 3 shows the FTIR spectra of MSP and MRL. As 340 f3 the treatment temperatures were increased, the bands assigned 341

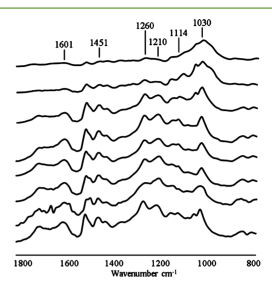


Figure 3. FTIR spectra of MSP and isolated lignin. From top to bottom: MSP, 170/180/190/200/210 °C MRL, KL, and 190 °C AL.

as aromatic skeleton $(1601/1508/1451/1424 \text{ cm}^{-1})^{25-28}$ were 342 strengthened significantly. These strong peaks suggested high 343 aromaticity of the residues after treatment. The peak at 1030- 344 1060 cm⁻¹ was assigned as C-O stretching of primary 345 alcohol. 25,27,28 It weakened as temperature rose, indicating a 346 better removal of polysaccharide at high temperature. The 347 overall trend of the FTIR spectra demonstrated that temper- 348 ature acts as an important factor in lignin isolation. At 349 treatment temperatures higher than 190 °C, the spectra of 350 MRL were very similar to that of KL. From 190 to 210 °C, the 351 peaks at 1114 cm⁻¹ (secondary alcohol)²⁹ and 1030 cm⁻¹ were 352 further weakened slightly. This may suggest that 210 °C MRL 353 was purer than 190 °C MRL. However, as shown in Table 1, 354 the 190 °C treatment rendered 18 wt % of the lignin acid 355 soluble. Therefore, an isolation at 210 °C would solubilize more 356 lignin, result in lower lignin yield, and perhaps trigger further 357 structural changes away from native lignin. Furthermore, the 358 tube pressure of the 210 °C experiment was 100 psi higher than 359 that of 190 °C (Figure S3). Therefore, due to lignin yield and 360 safety reasons, 190 °C seemed a suitable temperature for this 361 current protocol.

The FTIR spectra of 190 °C MRL and 190 °C AL showed 363 similar general trends. However, the peak at 1260 cm⁻¹ was 364 stronger in 190 °C AL than that in 190 °C MRL. This peak 365 could be ascribed to ether bonds, especially alkyl aryl ethers.²⁵ 366

SSNMR. Figure 4 shows the spectra of SSNMR spectra of 367 f4 MSP and various isolated lignin samples. The peak at 55 ppm 368 was as being attributable to methoxyl carbons. ^{29,30} This peak 369 was strengthened in isolated lignin samples, because the 370 monomer of softwood lignin, the guaiacyl unit (G-unit), 371 contains one methoxyl side chain. Comparing the spectra of 372 MSP and 170/190 °C MRL, it was obvious that the peaks 373 between 109 and 162 ppm were much stronger after microwave 374 treatment. According to Mao et al., ²⁹ the peaks in the range 375 between 108 and 60 ppm can be attributed to aliphatic carbons 376

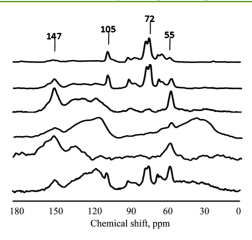


Figure 4. SSNMR spectra of MSP and isolated lignin. From top to bottom: MSP, 170 $^{\circ}$ C MRL, 190 $^{\circ}$ C MRL, KL, KL (CPNQS), and 190 $^{\circ}$ C AL.

mainly from carbohydrates and side chains of lignin, such as the peaks at 72 and 105 ppm characteristic of C2, C3, C5, and C1 rates of carbons of cellulose, while peaks between 162 and 109 ppm were attributed to carbon atoms in benzenoid rings that provided strong evidence for the existence of lignin in their samples. The major peaks in this zone were located at 147 ppm (aromatic C-O), and 125 ppm (hydrogen-bearing aromatic carbon not most adjacent to oxygen functionalities), and 114 ppm (aromatic carbon ortho to phenolic C-OH moieties).

The spectra of 190 °C MRL and KL showed significant differences. There were two wide bands at 30-50 ppm and 120–135 ppm for KL spectrum. Research^{29,33,34} showed that these two bands can be attributed to CH2 carbons and CH carbons, respectively. When processing the KL spectrum using the CPNQS methodology that suppresses the CH₂/CH band, the spectrum became similar to that of 190 °C MRL. These data suggested that microwave isolation can keep the aromatic part of lignin intact; however, it appears to remove the aliphatic part to some extent. The monomers of lignin are phenylpropanoid in structure. They are based on a C₆-C₃ structure 398 that contains both aliphatic and aromatic carbons. Compared 399 with the aromatic C₆ moieties, the C₃ aliphatic side chains of 400 lignins are characterized by higher polarities, having higher O/ 401 C ratios than the aromatic parts of the structure. Microwaves 402 are more efficient in heating polar compounds and functional 403 groups, 12 so the side chain is more likely to be modified or 404 cleaved during lignin isolation. As a result, lignin isolated using 405 microwave heating has a higher proportion of intact aromatic 406 rings and a lower proportion of intact side chains than does the

lignin isolated using conventional heating at the same 407 temperature. This fact will benefit the application of isolated 408 lignin as a potential source for production of low molecular 409 weight aromatic compounds.

The SSNMR spectrum of 190 °C AL showed a strong peak $_{411}$ at 72 ppm, suggesting severe sugar contamination. Similar to $_{412}$ the spectrum of KL (normal CP spectrum), there was a band at $_{413}$ 120–135 ppm, suggesting a high content of CH $_{2}$ group in $_{414}$ lignin isolated by conventional acidolysis. These data add $_{415}$ further evidence to the hypothesis that MRL has a $_{416}$ proportionately higher aromatic carbon content and that $_{417}$ microwave heating at 190 °C results in significant cleavage of $_{418}$ the side chains of this type of lignin.

Py-GC/MS. MSP and the isolated lignin samples (190 °C 420 MRL, KL, and 190 °C AL) were analyzed by Py-GC/MS. From 421 the changes of peak area % of typical pyrolytic products, 422 especially phenolic compounds, the structure change and 423 degradation extent during lignin isolation can be investigated. 424 Because most polysaccharide had been removed, the phenolic 425 compounds were dominant in pyrolytic products of the three 426 lignin samples, while there were more pyrolytic products from 427 cellulose and hemicellulose in MSP, such as 2-propanone, 1- 428 hydroxy-/furfural/cyclopentane-1,2-dione. In Table 2, nine of 429 t2 the compounds identified in highest proportions from the Py- 430 GC/MS are listed together with their measured ion current 431 peak areas. In Table 1 it was shown that the lignin content of 432 190 °C MRL (80.64 wt %, dry basis) was 2.67 times that of 433 MSP (30.37 wt %, dry basis). When the ratios between the ion 434 current peak areas for the 190 °C MRL and those for MSP are 435 compared as shown in Figure 5a, it is apparent that the trend 436 f5 line has a slope of 2.99, which is in acceptable agreement with 437 the expected ratio of 2.67, suggesting that lignin was well- 438 preserved without significant degradation. Notably, two of the 439 nine compounds were significant outliers from the trend line, 2-440 methoxy-4-vinylphenol and (E)-isoeugenol. There are two $_{441}$ possible reasons that can explain why these two compounds do 442 not conform to the expected trend: (1) The precursors for 443 these compounds are concentrated around the periphery of the 444 3D lignin structure and are bonded covalently to carbohydrates 445 as part of the LCC, resulting in chemical modification of the 446 alkene groups during acid hydrolysis. (2) The compounds are 447 more or less evenly distributed through the 3D structure of the 448 lignin and do not survive the acidic conditions at 190 °C for 449 reasons that cannot be explained at present. The fact that 450 compound numbers 6 and 7 in Table 2 also contain double 451 bonds in the side chain and do fit closer to the trend line may 452 be seen as evidence favoring the former explanation. The trend 453 line between KL and MSP (Figure 5b) was also somewhat 454 lower than that in Figure 5a, showing that there were 455

Table 2. Comparisons of Phenolic Compounds Peak Area (%) of MSP and Isolated Lignin

no.	compounds	MSP	190 °C MRL	KL
1	1,2-benzenediol, 4-methyl-	0.42	2.51	2.62
2	phenol, 4-ethyl-2-methoxy-	1.20	3.66	3.70
3	creosol	6.20	18.67	13.90
4	phenol, 2-methoxy-	3.17	8.45	9.73
5	vanillin	1.36	2.67	1.29
6	phenol, 2-methoxy-5-(1-propenyl)-, (E)-	0.99	1.87	1.43
7	phenol, 2-methoxy-4-(1-propenyl)-, (Z) -	0.39	0.65	0.31
8	2-methoxy-4-vinylphenol	4.04	4.92	3.58
9	trans-isoeugenol	3.78	3.20	1.37

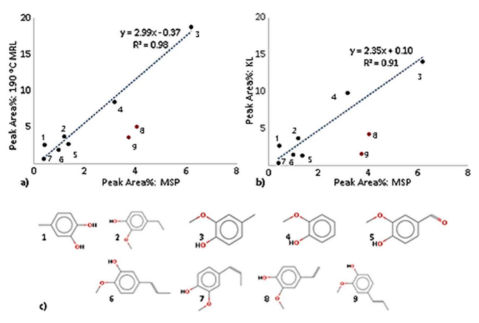


Figure 5. Peak area % of phenolic compounds according py-GC/MS analysis. (a) MSP vs 190 °C MRL; (b) MSP vs KL. (c) Compound structures.

456 proportionately fewer aromatic compounds in pyrolytic 457 products of KL than that of 190 °C MRL. This was probably 458 becasue there were more aliphatic compounds in KL due to less 459 side chain modification than that in 190 °C MRL, which is 460 consistent with the results of SSNMR analysis presented above. When the volatile products produced by Py-GC/MC of 190 462 °C AL were compared with those obtained from 190 °C MRL, 463 it was evident that pyrolysis products derived from carbohy-464 drates, such as 5-hydroxymethyl furfural (0.60% in 190 °C AL, 465 0.22% in 190 °C MRL) and D-allose (2.06% in 190 °C AL, 466 undetectable in 190 °C MRL), were evident with higher peak 467 areas in the case of 190 °C AL. An interesting fact was that one of main pyrolytic products, creosol, showed a higher peak area 469 % in 190 °C AL (25.0%) than that in 190 °C MRL (18.7%), 470 though the latter was purer lignin and less contaminated with carbohydrates. Fleck³⁵ observed that some model lignin dimers, such as conidendrin and di-isoeugenol in which the two 473 monomers are linked by a saturated ring, did not produce creosol during pyrolysis. Fleck found that certain interlinkages, 475 such as an indane ring, could effectively prevent the formation 476 of creosol under pyrolytic conditions. Furthermore, Fleck³⁵ pointed out that creosol was one of the main pyrolytic products 478 of coniferin which is a glucoside of coniferyl alcohol, so sugar 479 contamination actually could increase the yield of creosol to 480 some extent. It is arguable that these two factors explain why 481 190 °C AL with the higher carbohydrate content produced 482 more creosol. It is also possible that some of the structural 483 changes in the side chains of the lignin promoted by microwave 484 heating lead to formation of new cyclic aliphatic interlinkages 485 between monomeric units that are in close proximity within the 486 3D stucture and that these changes also serve to reduce creosol 487 yields from Py-GC/MS of 190 °C MRL.

488 CONCLUSIONS

489 It has been demonstrated that a pure form of lignin relatively 490 uncontaminated by residual carbohydrates can be produced 491 rapidly and efficiently by brief (10 min) microwave heating of 492 mixed softwood pellets (MSP) at 190 °C in dilute aqueous 493 sulfuric acid. The type of lignin produced by this new method,

designated as 190 °C MRL, has both higher yield and purity 494 than equivalent material produced by conventional heating to 495 190 °C in aqueous sulfuric acid at the same concentration in an 496 autoclave for the same time. The latter material has been 497 designated 190 °C AL (acidolysis lignin). It has been shown 498 that 190 °C MRL is of high aromaticity due to the modification 499 of lignin side chains. The Py-GC/MS results from the two 500 types of lignin indicate that some formation of cyclic aliphatic 501 linkage occurs between the side chains of monomeric units that 502 are in close proximity when microwave heating at 190 °C is 503 applied. The techniques applied using comparative Py-GC/MS 504 on lignin samples obtained by differing techniques have general 505 application in identifying structural changes occurring during 506 lignin isolation.

In general, the research results show that high-temperature 508 microwave treatment is a powerful tool for lignin isolation. 509 High efficiency, a simple protocol, and high lignin yield are its 510 most significant advantages. It is potentially a very promising 511 method for high-quality lignin preparation.

ASSOCIATED CONTENT

S Supporting Information

The Supporting Information is available free of charge on the 515 ACS Publications website at DOI: 10.1021/acssusche-516 meng.6b02545.

Element and ICP analysis of feedstock; TAPPI T222 ₅₁₈ method; GC/MS spectra and compounds lists; pressure ₅₁₉ and temperature comparisons during experiment (PDF) ₅₂₀

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531 Notes

532 The authors declare no competing financial interest.

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REFERENCES

- 540 (1) Méchin, V.; Baumberger, S.; Pollet, B.; Lapierre, C. Peroxidase 541 activity can dictate the in vitro lignin dehydrogenative polymer 542 structure. *Phytochemistry* **2007**, *68* (4), 571–579.
- 543 (2) Alonso, M. V.; Oliet, M.; Garcia, J.; Rodriguez, F.; Echeverría, J. 544 Gelation and isoconversional kinetic analysis of lignin-phenol—545 formaldehyde resol resins cure. *Chem. Eng. J.* **2006**, *122* (3), 159–166.
- 546 (3) Wahyudiono; Sasaki, M.; Goto, M. Recovery of phenolic 547 compounds through the decomposition of lignin in near and
- 548 supercritical water. *Chem. Eng. Process.* **2008**, 47 (9-10), 1609–1619. 549 (4) Yuan, T. Q.; Xu, F.; Sun, R. C. Role of lignin in a biorefinery:
- 550 separation characterization and valorization. *J. Chem. Technol.* 551 *Biotechnol.* **2013**, 88 (3), 346–352.
- 552 (5) Tuomela, M.; Vikman, M.; Hatakka, A.; Itävaara, M. 553 Biodegradation of lignin in a compost environment: a review. 554 *Bioresour. Technol.* **2000**, *72* (2), 169–183.
- 555 (6) Buranov, A. U.; Mazza, G. Lignin in straw of herbaceous crops. 556 Ind. Crops Prod. 2008, 28 (3), 237–259.
- 557 (7) Kubo, S.; Uraki, Y.; Sano, Y. Preparation of carbon fibers from 558 softwood lignin by atmospheric acetic acid pulping. *Carbon* **1998**, *36* 559 (7), 1119–1124.
- (8) Ragauskas, A. J.; Beckham, G. T.; Biddy, M. J.; Chandra, R.; 661 Chen, F.; Davis, M. F.; Davison, B. H.; Dixon, R. A.; Gilna, P.; Keller, 662 M.; et al. Lignin valorization: improving lignin processing in the 663 biorefinery. *Science* **2014**, 344 (6185), 1246843.
- 564 (9) Acid-Insoluble Lignin in Wood and Pulp; test method T222 om-02; 565 TAPPI: Peachtree Corners, GA, 2002.
- 566 (10) Fujimoto, A.; Matsumoto, Y.; Chang, H. M.; Meshitsuka, G. 567 Quantitative evaluation of milling effects on lignin structure during the 568 isolation process of milled wood lignin. *J. Wood Sci.* **2005**, *51* (1), 89–569 91.
- 570 (11) Wu, S.; Argyropoulos, D. S. An improved method for isolating 571 lignin in high yield and purity. *J. Pulp Pap. Sci.* **2003**, 29 (7), 235–240.
- 572 (12) Fan, J.; De Bruyn, M.; Budarin, V. L.; Gronnow, M. J.; 573 Shuttleworth, P. S.; Breeden, S.; Macquarrie, D. J.; Clark, J. H. Direct 574 microwave-assisted hydrothermal depolymerization of cellulose. *J. Am.*
- 575 Chem. Soc. **2013**, 135 (32), 11728–11731.
- 576 (13) Gulbrandsen, T. A.; Johnsen, I. A.; Opedal, M. T.; Toven, K.; 577 Øyaas, K.; Pranovich, A.; Mikkola, J. P.; Hoff, B. H. Extracting 578 hemicelluloses from softwood and bagasse as oligosaccharides using 579 pure water and microwave heating. *Cell. Chem. Tech.* **2015**, *49* (2), 580 117–126.
- 581 (14) Borges, F. C.; Du, Z.; Xie, Q.; Trierweiler, J. O.; Cheng, Y.; 582 Wan, Y.; Liu, Y.; Zhu, R.; Lin, X.; Chen, P.; Ruan, R. Fast microwave 583 assisted pyrolysis of biomass using microwave absorbent. *Bioresour*. 584 *Technol.* **2014**, *156*, 267–274.
- 585 (15) Zhou, S.; Liu, L.; Wang, B.; Xu, F.; Sun, R. Microwave-enhanced 586 extraction of lignin from birch in formic acid: Structural character-587 ization and antioxidant activity study. *Process Biochem.* **2012**, 47 (12), 588 1799–1806.
- 589 (16) Li, M. F.; Sun, S. N.; Xu, F.; Sun, R. C. Microwave-assisted 590 organic acid extraction of lignin from bamboo: Structure and 591 antioxidant activity investigation. *Food Chem.* **2012**, *134* (3), 1392–592 1398.

- (17) Zoia, L.; Orlandi, M.; Argyropoulos, D. S. Microwave-assisted 593 lignin isolation using the enzymatic mild acidolysis (EMAL) protocol. 594 *J. Agric. Food Chem.* **2008**, 56 (21), 10115–10122.
- (18) Gindl-Altmutter, W.; Obersriebnig, M.; Veigel, S.; Liebner, F. 596 Compatibility between cellulose and hydrophobic polymer provided 597 by microfibrillated lignocellulose. *ChemSusChem* **2015**, 8 (1), 87–91. 598
- (19) Rojo, E.; Peresin, M. S.; Sampson, W. W.; Hoeger, I. C.; 599 Vartiainen, J.; Laine, J.; Rojas, O. J. Comprehensive elucidation of the 600 effect of residual lignin on the physical, barrier, mechanical and surface 601 properties of nanocellulose films. *Green Chem.* **2015**, *17* (3), 1853–602
- (20) Argyropoulos, D. S.; Sun, Y.; Palus, E. Isolation of residual kraft 604 lignin in high yield and purity. *J. Pulp Pap. Sci.* **2002**, 28 (2), 50–54. 605 (21) Adler, E. Lignin chemistry—past, present and future. *Wood Sci.* 606
- Technol. 1977, 11 (3), 169–218.
- (22) Lin, J.; Kubo, S.; Yamada, T.; Koda, K.; Uraki, Y. Chemical 608 thermostabilization for the preparation of carbon fibers from softwood 609 lignin. *BioResources* **2012**, *7* (4), 5634–5646.
- (23) Wang, G.; Li, W.; Li, B.; Chen, H. TG study on pyrolysis of 611 biomass and its three components under syngas. *Fuel* **2008**, 87 (4), 612 552–558.
- (24) Biagini, E.; Barontini, F.; Tognotti, L. Devolatilization of 614 biomass fuels and biomass components studied by TG/FTIR 615 technique. *Ind. Eng. Chem. Res.* **2006**, 45 (13), 4486–4493.
- (25) Degen, I. A. Tables of Characteristic Group Frequencies for the 617 Interpretation of Infrared and Raman Spectra; Acolyte: Harrow, U.K., 618 1997
- (26) Chen, J. Y.; Shimizu, Y.; Takai, M.; Hayashi, J. A method for 620 isolation of milled-wood lignin involving solvent swelling prior to 621 enzyme treatment. *Wood Sci. Technol.* **1995**, 29 (4), 295–306.
- (27) Huang, Y.; Wang, L.; Chao, Y.; Nawawi, D. S.; Akiyama, T.; 623 Yokoyama, T.; Matsumoto, Y. Analysis of lignin aromatic structure in 624 wood based on the IR spectrum. *J. Wood Chem. Technol.* **2012**, 32 (4), 625 294–303.
- (28) Kline, L. M.; Hayes, D. G.; Womac, A. R.; Labbe, N. Simplified 627 determination of lignin content in hard and soft woods via UV-628 spectrophotometric analysis of biomass dissolved in ionic liquids. 629 *BioResour.* **2010**, *5* (3), 1366–1383.
- (29) Mao, J.; Holtman, K. M.; Scott, J. T.; Kadla, J. F.; Schmidt-Rohr, 631 K. Differences between lignin in unprocessed wood, milled wood, 632 mutant wood, and extracted lignin detected by 13C solid-state NMR. *J.* 633 Agric. Food Chem. **2006**, 54 (26), 9677–9686.
- (30) Bardet, M.; Foray, M. F.; Trân, Q. K. High-resolution solid-state 635 CPMAS NMR study of archaeological woods. *Anal. Chem.* **2002**, 74 636 (17), 4386–4390.
- (31) Hatfield, G. R.; Maciel, G. E.; Erbatur, O.; Erbatur, G. 638 Qualitative and quantitative analysis of solid lignin samples by carbon- 639 13 nuclear magnetic resonance spectrometry. *Anal. Chem.* **1987**, 59 640 (1), 172–179.
- (32) Dick-Perez, M.; Wang, T.; Salazar, A.; Zabotina, O. A.; Hong, 642 M. Multidimensional solid-state NMR studies of the structure and 643 dynamics of pectic polysaccharides in uniformly 13C-labeled 644 Arabidopsis primary cell walls. *Magn. Reson. Chem.* **2012**, *50* (8), 645 539–550.
- (33) Martinez, A. T.; Almendros, G.; González-Vila, F. J.; Fründ, R. 647 Solid-state spectroscopic analysis of lignins from several Austral 648 hardwoods. *Solid State Nucl. Magn. Reson.* **1999**, *15* (1), 41–48.
- (34) Nogueira, R. F.; Boffo, E. F.; Tavares, M. I. B.; Moreira, L. A.; 650 Tavares, L. A.; Ferreira, A. G. The Use of Solid State NMR to Evaluate 651 the Carbohydrates in Commercial Coffee Granules. *Food Nutr. Sci.* 652 **2011**, 2 (4), 350.
- (35) Fleck, J. A. The investigation of peracetic acid-oxidized loblolly 654 pine by pyrolysis-gas chromatography-mass spectrometry. Ph.D. 655 Thesis. Lawrence University, Appleton, WI, 1975.